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#### **Published**

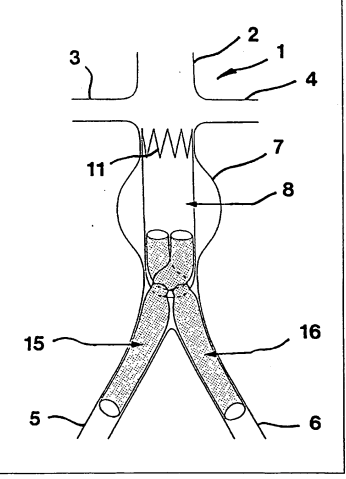
With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: AN ENDOVASCULAR GRAFT PROSTHESIS AND AN IMPLANTATION METHOD FOR SUCH A PROSTHESIS

#### (57) Abstract

An endovascular graft prosthesis for arrangement at or in the vicinity of a bifurcation in the arterial system of a patient, e.g. for purpose of repairing an aneurysm or the aortic bifurcation comprises a substantially tubular main body for location in the principal upstream arteria above the bifurcation such as the aorta and substantially tubular legs joining said main body in a bifurcation and extending into each of two branch arteries such as the iliac arteries. The main body (8) is substantially bag-shaped with an open proximal upstream end and a distal downstream bottom region in which two outlet openings (9, 10) are provided and is attachable to the inner side of the principal arteria (2) by means of an expandable stent device (11). The legs are made as separate expandable leg stent devices (15, 16) which may be introduced in a collapsed condition through the branch arteries and into the outlet openings (9, 10) of the main body by means of guide wires (13, 14) to engage against the rim of the corresponding outlet opening to provide a leakage-free bifurcated graft prosthesis.



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An endovascular graft prosthesis and an implantation method for such a prosthesis.

The invention relates to an endovascular graft prosthesis for arrangement at or in the vicinity of a 5 bifurcation in the arterial system of a patient and comprising a substantially tubular main body for location in an upstream arteria above the bifurcation and substantially tubular legs joining said main body and extending via the bifurcation into each of two 10 downstream branch arteries, said main body being made of a flexible microporous and surgically implantable woven material unpenetratable to blood.

In particular, the invention is concerned with the repair of an aneurysm in the vicinity of the aortic 15 bifurcation, but it may also be applied to other parts of the arterial system where a principal upstream arteria bifurcates into two branch arteries.

In order to prevent an aortic aneurysm, in particular in the lower part of the aorta close to the 20 aortic bifurcation from causing a dangerous rupture of the aortic wall it is known to deploy a graft prosthesis in the region of the vessel affected by such an aneurysm.

An aortic aneurysm may develop as a result of a reduction of the strength of the aortic wall whereby the diameter of the affected part of the aorta may increase to more than 5 cm. Such expansions may result in flow irregularities and promotion of deposits of coagulated blood in the affected region. At increased expansion the remaining strength of the aortic wall will naturally decrease and may ultimately result in rupture of the vessel with an inherent danger of acute bleeding.

With conventional prior art endovascular prosthesis for aortic implantation surgical opening of

the actual vascular section will be necessary for deployment of the prosthesis. For this purpose a partial cut is made in the wall of the aneurysm to introduce the prosthesis formed as an integral unit of a plastic 5 material and secure it by sewing.

From EP-A-0508473 and EP-A-0539237 bifurcated graft prosthesis are known which may be transluminarly implanted for the repair of an aneurysm at or in the vicinity of the aortic bifurcation. In both cases the 10 bifurcated prosthesis is made as an integral unit with a main body and two tubular legs joining the main body in a bifurcation. Due to this design the implantation operation becomes relatively complicated since the integral unit must be introduced through one of the 15 iliac arteries with one of the legs in a fold-over condition until the graft is disposed proximal of the aortic bifurcation following which the proximal extremity of the prosthesis must be secured upstream of the actual vascular section and the folded overleg must 20 be pulled down into the other iliac arteria.

It is the object of the invention to provide an endovascular graft prosthesis of the kind set forth for transluminal implantation at the aortic bifurcation by a considerably simpler implantation operation than the above-mentioned prior art solutions.

In order to achieve this an endovascular graft prosthesis according to the invention is characterized in that the main body is substantially bag-shaped with an open proximal upstream end and a distal downstream bottom region in which two outlet openings are provided, said main body being radially expandable and attachable in a radially expanded condition to the inner side of said upstream arteria upstream of the bifurcation by fixation means, said legs being made as separate resilient and radially expandable leg stent devices

adapted for introduction in a collapsed condition through said branch arteries and into said outlet openings, each of said stent devices being engageable in its radially expanded condition against the rim of the corresponding outlet opening to provide a leakage-free bifurcated graft prosthesis.

By making up the prosthesis from a number of separate components which may be sequentially introduced in the arterial system by percutaneous operations through small openings with a punctual diameter up to 5 mm the components may be endovascularly assembled to a complete prosthesis after deployment in the actual vascular section. Thereby, the prosthesis according to the invention may also be applied for repair of an aneurysm extending into the iliac arteries.

The invention further relates to a method for implanting an endovascular graft prosthesis for deployment at or in the vicinity, a bifurcation in the arterial system of a patient and associated branch 20 arteries.

the invention this According to characterized by the steps of introducing through a first branch arteria in an upstream direction by means of a first guide wire a separate, radially expandable 25 and substantially bag shaped main body into an upstream arteria to extend into a region thereof above the bifurcation, said main body having an open proximal upstream end with associated fixation means and a distal downstream bottom region in which two outlet openings 30 are provided, expanding said main body radially in said upstream arteria with said proximal end attached to the wall thereof, and introducing by means of said further quide wires a radially expandable leg stent device through each of the branch arteries into each of said 35 outlet openings.

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By this method the main body consisting of a micro porous woven material unpenetratable to blood may be guided by means of a guide wire from a puncture in an iliac arteria into the aorta and deployed there with a 5 downstream orientation of the distal bottom region of the body with respect to the blood flow direction. Thereby, the guide wire will be coaxially located in one or both of the outlet openings of the bag-shaped body. In spite of the two outlet openings the main body will 10 be sufficiently expanded by the blood flow in its bottom region. In this condition, the main body is secured to the inner wall of the aorta upstream of the bifurcation by means of a radially expandable stent device which may typically be a self-expandable metallic stent. Following 15 subsequent introduction of guide wires through the two outlet openings and across the lumen of the stent device a leg stent device may be introduced over each guide wire to extend at least through the distance between the bottom region of the main body and the junction of the 20 iliac arteries. The leg stent devices forming separate components of the prosthesis may have an elastic covering over their entire length and are also radially expandable.

The introduction of the two leg stent devices
25 which may also be self-expandable metallic stents may
take place by use of conventional catheter technique
whereby the guide wire and the catheter are guided from
one outlet opening through the other and into the
opposite iliac arteria from where it may be guided in
30 a conventional way through the vessel to the skin. By
drawing of the curved guide wire extending through the
outlet openings the position of the main body may still
be corrected. Subsequently, two catheters may be
introduced into the outlet openings over the guide wire
35 thus positioned from the two groins to allow coaxial

introduction of two guide wires to replace the two catheters. Subsequently the two leg stents covered by woven material are introduced over the positioned guide wires which can take place simultaneously for both stent devices.

In the expanded condition the two leg stents will press against the rim of the outlet openings and complete the main body into a leakage free endovascular prosthesis with a bifurcation secured in the aorta.

Preferably the main stent is attached to the bagshaped main body with the downstream distal end overlapped by the upstream proximal end of the main body.

By a further development of the invention the 15 endovascular deployment of the prosthesis may be substantially facilitated if the main stent device and the leg stent devices in a manner known per se are formed by self-expandable metal stents.

According to a further embodiment of the invention 20 the rim of each outlet opening may diverge in the downstream direction.

By this measure there will be formed in each outlet opening close to the inner side of the bottom region of the bag-shaped main body a radially inwardly projecting opening rim against which the external side of the leg stent may engage tightly whereby the security against leakage will be increased.

Further, according to an embodiment of the invention the maximum external diameter of each of the 30 leg stents may be 2-4 mm greater than the minimum rim diameter of each outlet opening.

Thereby, the components may be connected in a sufficiently stable and leakage free manner without any requirement for additional means for this purpose.

35 The invention will be further explained with

reference to an embodiment shown on the schematical drawing in which

fig. 1 shows a bag-shaped main body of a prosthesis according to the invention deployed in 5 expanded condition within an aortic aneurysm,

fig. 2 illustrates the introduction of two guide wires into the main body of fig. 1 and

fig. 3 the main body completed with two leg stents into a prosthesis with a bifurcation.

As example of the application of the invention, figures 1 to 3 illustrate schematically an arterial system 1 in which the abdominal part of the aorta 2 extending between two branch arteries 3,4 leading to the kidneys and a bifurcation joining two iliac arteries 5,6 has been damaged by a balloon shaped aneurysm 7.

As illustrated in figure 1 a bag-shaped main body made of a micro-porous woven material unpenetratable to blood is first introduced in the region of the aneurysm 7 in the aorta 2, said main body being formed in a 20 bottom region with two outlet openings 9,10. The main body 8 is provided with a self-expandable metallic stent 11 and is introduced in contracted condition through an iliac arteria 5 over a guide wire 12 and deployed in the aorta 2 in such a way that the metallic stent 11 will 25 press at least the upper edge of the main body 8 tightly against the inner side of the aorta 2 above the aneurysm 7. In the bottom region of the main body 8 the guide wire 12 may as shown in dashed lines in fig. 1 be bent so as to allow it to be guided outwards through the 30 outlet opening and the other iliac arteria 6. Thereby a catheter not shown may be introduced over each of the free ends of the guide wire 12 until the main body 8 which has expanded due to the action of the blood flow. After removal of the guide wire 12 a single guide wire 35 13,14 may be introduced through each catheter to

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finalize the deployment preparation as illustrated in figure 2.

Over each of the guide wires 13,14 a self-expandable metallic stent 15,16 with an elastic covering 5 may now be introduced in a conventional way. As illustrated in fig. 3 a proximal end part of each of the metallic stents 15,16 projects into the interior of the main body whereas a distal end part of each metallic stent projects relatively deep into the iliac arteria 10 5,6.

In the expanded condition illustrated in fig. 3 the metallic stents 15,16 are contracted in the rim region of each of outlet openings 9 and 10 and will thereby be firmly and tightly connected with the main body 8. Thereby, the main body together with the leg stents 15 and 16 will form an endovascular aortic prosthesis overlapping the aneurysm 7 and being implantable by relatively simple percutaneous operations.

#### PATENT CLAIMS

- An endovascular graft prosthesis for arrangement at or in the vicinity of a bifurcation in the arterial system of a patient and comprising a substantially 5 tubular main body for location in an upstream arteria above the bifurcation and substantially tubular legs joining said main body and extending via the bifurcation into each of two downstream branch arteries, said main body being made of a flexible microporous and surgically 10 implantable woven material unpenetratable to blood, characterized in that the main body (8) is substantially bag-shaped with an open proximal upstream end and a distal downstream bottom region in which two outlet openings (9, 10) are provided, said main body being 15 radially expandable and attachable in a radially expanded condition to the inner side of said upstream arteria (2) upstream of the bifurcation by fixation means (11), said legs being made as separate resilient and radially expandable leg stent devices (15, 16) 20 adapted for introduction in a collapsed condition through said branch arteries and into said outlet openings (9,10), each of said stent devices (15, 16) being engageable in its radially expanded condition against the rim of the corresponding outlet opening to 25 provide a leakage-free bifurcated graft prosthesis.
- 2. An endovascular graft prosthesis as claimed in claim 1, characterized in that said fixation means comprises a main stent device (11) is attached to the bag-shaped main body (8) with a downstream distal end 30 of the main stent device (11) overlapped by the upstream proximal end of the main body (8).
- An endovascular graft prosthesis as claimed in claim 2, characterized in that said main stent device (11) and leg stent devices (15,16) are formed by 35 expandable metal stents.

- 4. An endovascular graft prosthesis as claimed in claim 1, 2 or 3, characterized in that the rim of each outlet opening (9, 10) diverges in the downstream direction.
- 5 5. An endovascular graft prosthesis as claimed in claim 4, characterized in that the maximum external diameter of each of the leg stents (15, 16) is 2-4 mm greater than the minimum rim diameter of each outlet opening (9, 10).
- 10 6. A method for implanting an endovascular graft prosthesis as claimed in any of the preceding claims for deployment at or in the vicinity, a bifurcation in the arterial system of a patient and associated branch arteries, characterized by the steps of
- introducing through a first branch arteria (5) in an upstream direction by means of a first guide wire (12) a separate, radially expandable and substantially bag shaped main body (8) into an upstream arteria to extend into a region thereof above the bifurcation, said main body having an open proximal upstream end with associated fixation means and a distal downstream bottom region in which two outlet openings (9, 10) are provided,

expanding said main body radially in said upstream 25 arteria with said proximal end attached to the wall thereof, and

introducing by means of said further guide wires (13, 14) a radially expandable leg stent device (15, 16) through each of the branch arteries (5, 6) into each of 30 said outlet openings.

7. An implantation method as claimed in claim 6, characterized in that a catheter is introduced through each of the branch arteries over each end of said first guide wire prior to the removal thereof, said catheters being subsequently used for introduction of said further

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guide wires.

Fig. 1 1/3

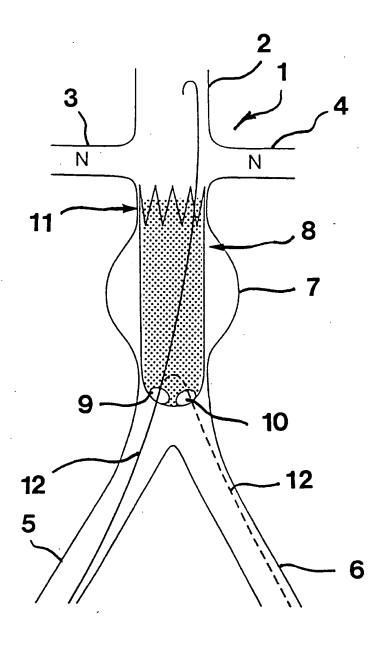


Fig. 2 2/3

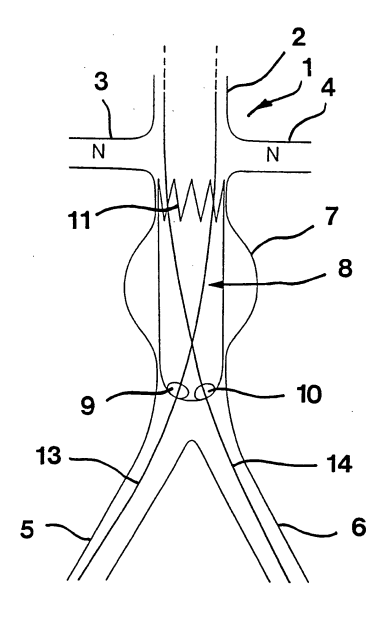
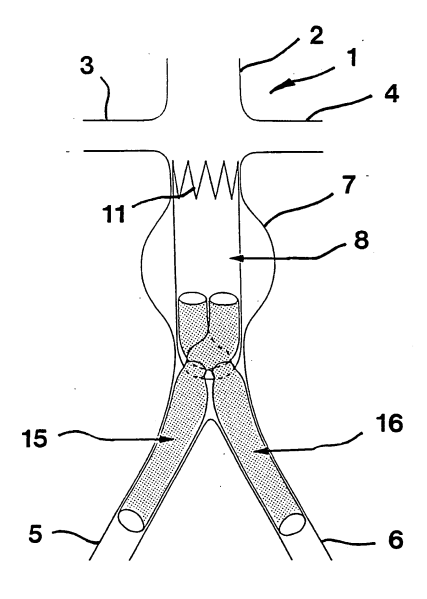


Fig. 3 3/3



International application No.

PCT/DK 94/00468

A. CLASS	SIFICATION OF SUBJECT MATTER		
IDCG. A	61E 2/06 // A61D 17/11		
According to	61F 2/06 // A61B 17/11 o International Patent Classification (IPC) or to both na	tional classification and IPC	
	S SEARCHED		
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IPC6: A	61B, A61F		
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DIALOUT	, EPODOC	•	
C. DOCU	MENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where app	propriate, of the relevant passages	Relevant to claim No.
A	EP, A1, 0461791 (BARONE, HECTOR I 18 December 1991 (18.12.91),	D. ET AL), figures 10-12	1-2
	<del></del> ,		
A	EP, A2, 0508473 (ENDOVASCULAR TE 14 October 1992 (14.10.92),		1-3
	***		
A .	EP, A1, 0539237 (COOK INCORPORATE (28.04.93), figure 47	ED), 28 April 1993	1-3
			•
A	EP, A1, 0551179 (EXPANDABLE GRAF 14 July 1993 (14.07.93), fig		1
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Furth	er documents are listed in the continuation of Box	C. X See patent family anne	x.
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Form PCT/ISA/210 (second sheet) (July 1992)

#### INTERNATIONAL SEARCH REPORT

International application No.

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Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)	
This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons	:
1. X Claims Nos.: 6-7 because they relate to subject matter not required to be searched by this Authority, namely:	
A method for treatment of the human or animal body by surgery or therapy (PCT, Rule 39.1 (iv))	
2. Claims Nos.:  because they relate to parts of the international application that do not comply with the prescribed requirements to suc an extent that no meaningful international search can be carried out, specifically:	ь
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a	).
Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)	$\neg$
This International Searching Authority found multiple inventions in this international application, as follows:	٦
	- 1
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1. As all required additional search fees were timely paid by the applicant, this international search report covers a searchable claims.	11
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payme of any additional fee.	nt
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:	п
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report restricted to the invention first mentioned in the claims; it is covered by claims Nos.:	is
Remark on Protest	
No protest accompanied the payment of additional search fees.	- 1

# INTERNATIONAL SEARCH REPORT

Information on patent family members

25/02/95

International application No. PCT/DK 94/00468

Patent document cited in search report		Publication date	Patent family member(s)		Publication date
EP-A1-	0461791	18/12/91	AU-B-	655418	22/12/94
			AU-A-	7754694	05/01/95
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